Abstract

Objectives—The metabolic changes in the brain of symptomatic subjects affected with Machado-Joseph disease have been previously documented using PET with fluorine-18-fluorodeoxyglucose (FDG). The aim of this study was to evaluate these changes in asymptomatic Machado-Joseph disease gene carriers.

Methods—Seven asymptomatic Machado-Joseph disease gene carriers, identified using a molecular test, and 10 normal control subjects were recruited for PET studies using FDG. Regional uptake ratios of FDG were calculated from the radioactivity of the cerebellar hemispheres, brainstem, and the temporal, parietal and occipital cortices, divided by the activity in the thalamus.

Results—In comparison with data obtained from normal control subjects, there was significantly decreased FDG utilisation in the cerebellar hemispheres, brainstem, and occipital cortex, and increased FDG metabolism in the parietal and temporal cortices of asymptomatic Machado-Joseph disease gene carriers, suggesting preclinical disease activity. Discriminant analysis of regional FDG uptake correctly classified genetic status (Machado-Joseph disease mutation carriers vs mutation negative subjects) in 25 of 25 subjects (100% sensitivity and 100% specificity), and clinical status (asymptomatic mutation carriers vs symptomatic patients) in 14 of 15 subjects (100% sensitivity and 85.7% specificity).

Conclusion—Subclinical changes of FDG consumption, as measured by non-invasive PET, can act as an objective marker of preclinical disease activity in Machado-Joseph disease.

Keywords: Machado-Joseph disease; positron emission tomography; asymptomatic gene carriers

Machado-Joseph disease (MIM 109150) is an autosomal dominantly inherited neurodegenerative disorder characterised pathologically by degeneration of the spinocerebellar tracts, dentate nuclei, pontine and vestibular nuclei, extrapyramidal structures (substantia nigra, locus coeruleus, and the pallidoluyian complex), and neuronal loss in motor cranial nerves, anterior horn cells, and the posterior root ganglion.1,2 The cerebral and cerebellar cortices and inferior olives are spared.3 Patients often present with cerebellar ataxia, pyramidal signs, and progressive external ophthalmoplegia. Bulging eyes, peripheral amyotrophy, and dystonia of varying severity may also be present.4 The natural course of this illness, with the average age at onset of symptoms in the late fourth decade of life, often results in many years of at risk status for those who have a parent with the disease. The insidious nature of Machado-Joseph disease often makes the exact age at onset of symptoms difficult to determine.

The Machado-Joseph disease gene has been isolated and characterised,5 and the mutation responsible for the disease has been shown to be an expansion of a trinucleotide CAG repeat that lies at the 3' terminal of the coding region.6 The identification of the characteristic CAG repeat expansion in the Machado-Joseph disease gene has since become the gold standard for the definitive diagnosis of the disease.

Positron emission tomography (PET) has been shown to be a useful tool in elucidating the pathophysiology of various movement disorders (Huntington's disease,4 Parkinson's disease,7 progressive supranuclear palsy,8 and spinocerebellar degeneration9–11). Using PET with fluorine-18-fluoro-2-deoxy-D-glucose (FDG), we showed that relative brain glucose metabolism was significantly diminished in the cerebellar hemispheres, cerebellar vermis, brainstem, and cerebral occipital cortex in patients affected with Machado-Joseph disease.11 The present study of asymptomatic Machado-Joseph disease gene carriers was undertaken to consider the question of whether regional brain hypometabolism can be found years before onset of disease, the sensitivity of the PET in the detection of asymptomatic Machado-Joseph disease gene carriers, and whether the length of CAG repeat correlates with the severity of these metabolic changes. In addition, we performed discriminant function analysis to assess whether the genetic and clinical status of the subjects could be predicted from brain functional values.

The study of metabolic changes in the brain of Machado-Joseph disease gene carriers is of importance as it could potentially identify subjects who may be suitable for therapeutic intervention in the near future to establish timing of treatment, to gauge response, and to halt or slow the disease process years before clinical presentation.

 Patients and methods

SUBJECTS

Previously, eight symptomatic subjects with Machado-Joseph disease were studied with PET using FDG.11 In this report seven
unrelated asymptomatic gene carriers of Machado-Joseph disease (four males and three females; age 27.7 (SD 11.4) years), identified by the presence of expanded CAG repeats (70.9 (SD 6.7)) in the Machado-Joseph disease gene, and 10 normal control subjects (four males and six females; age 46.9 (SD 11.8) years, range of CAG repeats 14–39) were recruited for investigation. All of the asymptomatic mutation carriers underwent careful neurological examination by an experienced board certified neurologist. None of them showed any sign of ataxia or other “soft” neurological signs. Informed consent was obtained from all subjects.

DNA ANALYSIS WITH QUANTITATIVE POLYMERASE CHAIN REACTION
Genomic DNA was isolated from buffy coat leucocytes as previously described, and analysed to determine the length of the CAG repeat in the Machado-Joseph disease gene. Polymerase chain reaction of genomic DNA was performed according to the method of Kawaguchi et al. Primers MJD52 and MJD25 were used to identify the triplet region. This technique identifies the number of CAG repeats in the Machado-Joseph disease gene.

PET STUDIES
All subjects were awake, taking no medication known to affect CNS function, and blindfolded during the examination. The imaging device was an eight ring Scanditronix PC4096–15WB whole body PET scanner (Scanditronix, Sweden) with an axial resolution of 6 mm and in plane resolution of 8 mm at the centre of the field of view. PET imaging was performed 45 to 75 minutes after injection of 10 mCi (370 MBq) FDG. Visual interpretation and semi-quantitative analysis of the PET images were conducted subsequently. The regions of interest (ROIs) were determined in the cerebellar hemispheres, brainstem, thalamus, and parietal, temporal, and occipital cortices by reference to an atlas of axial tomography. Extreme caution was exercised in the placement of ROIs to avoid potential contamination from adjacent anatomical structures. Data were collected from the ROIs by placing an 11×11 mm square over each thalamus, an 11×11 mm square over the occipital cortex, a 10×15 mm rectangle over the frontal, temporal, and parietal cortices (fig 1 C), a 11×22 mm parallelogram over each cerebellar hemisphere, and an 11×15 mm rectangle over the brainstem (fig 1 D). Each ROI was centred over a local peak in FDG metabolism. Uptake of FDG was calculated as nCi/ml. Regional uptake ratios were then calculated from the radioactivity of the cerebellar hemispheres, brainstem, and the temporal, parietal, and occipital cortices, divided by the activity in the thalamus.

Figure 1  FDG PET in subject 5 (A, B) who is an asymptomatic gene carrier of Machado-Joseph disease, and a normal control subject (C, D). Relative to the FDG uptake in the thalamus, the FDG uptake ratio is decreased in the cerebellar hemisphere, brainstem (B), and occipital cortex (A), but increased in the parietal and temporal cortices (A). Squares represent the regions of interest marked in the thalamus, occipital and frontal cortices, brainstem, and cerebellum. Colour bars indicate regional brain FDG uptake in nCi/ml extending from 0.0 to 200 (A, B) and 0.0 to 400 (C, D).

STATISTICAL ANALYSIS
Age, CAG repeat number, and regional brain FDG uptake ratios are presented as range and mean (SD). Differences of brain FDG uptake ratios in asymptomatic gene carriers of Machado-Joseph disease, symptomatic patients with Machado-Joseph disease, and control subjects are presented as 95% confidence intervals (95% CIs). A Mann-Whitney U test was used to assess the statistical significance. The association between the age of asymptomatic gene carriers, the size of the trinucleotide repeat sequence, and the regional brain uptake ratios of FDG were evaluated through Spearman’s correlation analyses. Discriminant function analysis was used to categorise subjects on the basis of their FDG values in the cerebellar hemispheres, brainstem, and occipital, temporal, and parietal cortices. Specificity (true negative/true negative+false positive) and sensitivity (true positive/true positive+false negative) were then calculated, based on discriminant analysis categorisation obtained from each of the above mentioned variables.

Results
Figure 1 shows typical brain FDG consumption scans in one of the asymptomatic Machado-Joseph disease gene carriers (subject 5) and one normal control subject. Table 1 shows the age, the CAG repeat number in the Machado-Joseph disease gene, and the individual regional brain uptake ratios of FDG of the seven asymptomatic Machado-Joseph disease gene carriers.

Significantly lower regional brain uptake ratios of FDG compared with that of normal subjects, were found over the cerebellar hemisphere (Machado-Joseph disease gene carriers 0.72–0.84, control 0.88 (SD 0.03)), brainstem (Machado-Joseph disease gene carriers 0.72–0.84, control 0.88 (SD 0.03)), and occipital cortices (Machado-Joseph disease gene carriers 0.72–0.84, control 0.88 (SD 0.03)).
carriers 0.59–0.77, control 0.78 (SD 0.08), and the occipital cortex (Machado-Joseph disease gene carriers 0.78–0.97, control 0.92 (SD 0.04) (table 2 and fig 2). By contrast, significantly higher regional brain uptake ratios of FDG were found over the parietal (Machado-Joseph disease gene carriers 0.97–1.20, control 0.96 (SD 0.03)) and temporal (Machado-Joseph disease gene carriers 0.92–1.11, control 0.90 (SD 0.04)) cortices (table 2 and fig 2). Comparison of the regional FDG uptake ratios between the asymptomatic gene carriers and the patients with Machado-Joseph disease also disclosed significant differences, in the parietal, temporal, and occipital cortices and the cerebellum (table 2).

No significant correlation was found between the size of CAG repeat and the regional brain uptake ratios of FDG. However, a tentative trend towards correlation seemed to be present between the age of the asymptomatic gene carriers and the regional FDG uptake ratio in the cerebellar hemisphere ($r=0.83$) and occipital cortex ($r=0.77$).

Discriminant function analysis was performed on the three groups (asymptomatic Machado-Joseph disease mutation carriers, mutation negative controls, and patients with Machado-Joseph disease) considering FDG uptake in the cerebellar hemispheres, brainstem, and occipital, parietal, and temporal cortices each as separate variables. The best categorisations between normal controls and asymptomatic Machado-Joseph disease mutation carriers were obtained with cerebellar FDG uptake ratios which showed an 86% sensitivity and a 100% specificity, and with temporal FDG uptake ratios which disclosed a 71.4% sensitivity and a 100% specificity. However, when considering the two dependent variables together, discriminant analysis classification was consistent with the genetic status in 17 of 17 subjects with a 100% sensitivity and a 100% specificity. The best categorisation between normal controls and symptomatic subjects was obtained with cerebellar FDG uptake ratios, which showed a 100% sensitivity and 100% specificity. Discriminant function analysis on FDG uptake ratios between asymptomatic mutant gene carriers and symptomatic patients was consistent with clinical status in 14 of 15 subjects with a 100% sensitivity and an 85.7% specificity in the parietal cortex. The one remaining person (subject

### Table 1 Regional brain uptake ratios of FDG in asymptomatic gene carriers of Machado-Joseph disease (MJD)

<table>
<thead>
<tr>
<th>Asymptomatic MJD gene carriers</th>
<th>Control subjects (mean (SD))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subject 1</td>
<td>Subject 2</td>
</tr>
<tr>
<td>Age (y)</td>
<td>13</td>
</tr>
<tr>
<td>CAG repeat number</td>
<td>80</td>
</tr>
<tr>
<td>CH/TH</td>
<td>0.72 (82)</td>
</tr>
<tr>
<td>BS/TH</td>
<td>0.73 (94)</td>
</tr>
<tr>
<td>O/TH</td>
<td>0.84 (91)</td>
</tr>
<tr>
<td>P/TH</td>
<td>1.0 (104)</td>
</tr>
<tr>
<td>T/TH</td>
<td>0.95 (106)</td>
</tr>
</tbody>
</table>

Numbers in parentheses for the asymptomatic MJD gene carriers represent percentage of each ratio compared with the mean of control subjects.

CH = cerebellar hemisphere; BS = brainstem; O = occipital; P = parietal; T = temporal; TH = thalamus.

Figure 2 Scatter diagrams of local brain metabolic ratio for glucose in asymptomatic gene carriers (A) of Machado-Joseph disease compared with normal control subjects (C) and symptomatic people (S) with Machado-Joseph disease. Each point represents the average value for each case in the structure specified. The triangles with error bars depict the mean (SD) value for each group. CH = cerebellar hemisphere; BS = brainstem; O = occipital; P = parietal; T = temporal; TH = thalamus. *p<0.05; **p<0.01; *** p<0.001.
Table 2  Comparison of regional brain uptake ratios of FDG in asymptomatic Machado–Joseph disease (MJD) gene carriers, patients with Machado–Joseph disease, and normal controls

<table>
<thead>
<tr>
<th>Patients with MJD (n=8)</th>
<th>Asymptomatic gene carriers (n=7)</th>
<th>Control subjects (n=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y, mean (SD))</td>
<td>28.9 (10.3)</td>
<td>27.7 (11.4)</td>
</tr>
<tr>
<td>CH/TH (0.73 (SD 0.05))</td>
<td>0.79 (SD 0.04)</td>
<td>0.79 (0.04)</td>
</tr>
<tr>
<td>BS/TH (0.63 (SD 0.07))</td>
<td>0.68 (SD 0.07)</td>
<td>0.68 (0.07)</td>
</tr>
<tr>
<td>O/TH (0.73 (SD 0.11))</td>
<td>0.86 (SD 0.07)</td>
<td>0.86 (0.07)</td>
</tr>
<tr>
<td>P/TH (0.91 (SD 0.01))</td>
<td>1.07 (SD 0.09)</td>
<td>1.07 (0.09)</td>
</tr>
<tr>
<td>T/TH (0.87 (SD 0.09))</td>
<td>1.02 (SD 0.07)</td>
<td>1.02 (0.07)</td>
</tr>
</tbody>
</table>

*p<0.05; **p<0.01; ***p<0.001
†Difference between regional FDG uptake ratios of subjects affected with MJD and that of control subjects.
‡Difference between regional FDG uptake ratios of subjects affected with MJD and that of asymptomatic gene carriers.
§Difference between regional FDG uptake ratios of asymptomatic gene carriers and those of control subjects.

CH = cerebellar hemisphere; BS = brainstem; O = occipital; P = parietal; T = temporal; TH = thalamus.

2) was an asymptomatic mutation carrier who was still serving in the army at the time of the PET measurement but was assigned to the category of symptomatic patients, indicating that this subject had an FDG uptake ratio in the parietal cortex indistinguishable from that in the affected patients.

Discussion

The role of PET in detecting subclinical disease has only been fully exploited in Huntington's disease.22 This is the first study to consider brain metabolism by PET in asymptomatic Machado-Joseph disease mutation carriers. Our results clearly show that subclinical changes in FDG consumption can be detected in vivo in all asymptomatic carriers of the Machado-Joseph disease mutation (table 1 and fig 2).

The difference in the age of asymptomatic gene carriers (27.7 (SD 11.4) years), symptomatic patients (28.9 (SD 10.3) years), and that of 10 normal control subjects (46.9 (SD 11.8) years) would not have any impact on baseline FDG concentrations (table 2).17 18 In an initial study of 21 subjects, later extended to 40 subjects, it was found that mean hemispheric cerebral metabolic rate for glucose, and regional metabolic rate for glucose in bilaterally symmetric and midline brain regions that were examined, were not correlated significantly with age.19 20

The thalamus was selected as the reference region for normalisation in our study for the following reasons: (1) The thalamus has been repeatedly shown to be unaffected and have a normal FDG metabolism in cases of olivopontocerebellar atrophy9 10 21 and Machado-Joseph disease.22 (2) FDG uptake could be calculated in the thalamus as well as in critical cerebral cortices from the same plane, allowing better comparison of data between these regions.

The metabolic changes detected with PET provide valuable information on the localisation of disease involvement. Using PET, Gilman et al found significant hypometabolism in the cerebellar hemispheres, cerebellar vermis, and brainstem in patients with sporadic and dominant olivopontocerebellar atrophy.9 10 Although Machado-Joseph disease has been regarded as a disease that severely affects the cerebellum and brainstem and largely spares the cerebrum, our previous in vivo PET study, aimed specifically at patients with Machado-Joseph disease, disclosed that relative brain metabolism was significantly diminished in the cerebral occipital cortex, as well as the cerebellar hemispheres, cerebellar vermis, and brainstem,11 which was consistent with the results of Taniwaki et al.22 The finding of regional changes of FDG consumption in asymptomatic gene carriers in this report indicates involvement of these structures preceding the onset of symptoms, further underlining the importance of our finding. Although atrophy could undoubtedly contribute to the hypometabolism through partial volume effects, no evidence of cerebellum or brainstem atrophy was found in two of our presymptomatic subjects who underwent cranial CT.

Our results strongly suggest that the brain metabolism is also regionally increased in the temporal and parietal areas early in the course of Machado-Joseph disease (table 2 and fig 2). This increased glucose metabolism was not seen in affected patients with Machado-Joseph disease,11 and could result from a metabolic defect that impairs energy metabolism, resulting in increased glycolysis.23 We suspect that the cortices in these regions may be affected initially, as indicated by the increase in the FDG uptake ratio in our asymptomatic Machado-Joseph disease gene carriers. However, these regions may be less vulnerable to the harmful effects of this metabolic defect, leading to the slower decline in FDG uptake ratio in symptomatic cases of Machado-Joseph disease. Alternatively, the failure to see such raised relative parietal and temporal metabolism in symptomatic cases of Machado-Joseph disease may reflect an increased cortical involvement of the disease in these later stage patients. These two viewpoints both seem to deserve consideration when the findings of our study are considered. There is also the remote possibility that this finding is an artefact from the normalisation procedure using the thalamus as a reference area or the use of an older control group with potentially reduced brain metabolism. However, this normalisation is unlikely to have generated such an artefact in our study because similar assessments have
PET evaluation of Machado-Joseph disease gene carriers

 However, there may be considerable variation of the same family in different generations (anticipation phenomenon), which makes estimation of the difference very unreliable.

Measurement of regional brain metabolism by PET is non-invasive, and subclinical changes of FDG consumption seem to be an objective marker of disease activity. PET studies may have an important part to play in the selection and monitoring of asymptomatic Machado-Joseph disease gene carriers in future therapeutic trials of neuroprotective agents, infusions of growth factors, or implantation of fetal cells. However, any increases in FDG consumption that were to follow the use of neuroprotective agents may inversely correlate with the duration of abnormal FDG uptake (table 1), despite the fact that a relatively high or low FDG uptake compared with a normal mean is not necessarily a good indication for neuronal dysfunction.

Previously we found that the CAG repeat in the Machado-Joseph disease gene was inversely correlated with the age at onset ($r=-0.77$) and the duration of the illness with the cortical hypometabolism ($r=-0.80$). With only seven asymptomatic Machado-Joseph disease gene carriers, covering a relatively narrow range of increased CAG repeat length, this study lacks the statistical power to show any correlation between the size of the CAG repeat sequence and the regional brain uptake ratios of FDG in the asymptomatic Machado-Joseph disease gene carriers. Moreover, although some linear trends towards a correlation could be seen between the age of the asymptomatic gene carriers and the regional FDG uptake ratios in the cerebellar hemisphere and the occipital cortex, these are also of questionable importance because of the few gene carriers studied.

The glucose metabolism data in the cerebellum, temporal, and parietal cortices were consistent with genetic status (Machado-Joseph disease mutation gene carriers vs mutation negative subjects) in 25 of 25 of subjects with a sensitivity of 100% and a specificity of 100%. Furthermore, the clinical status (asymptomatic vs clinically affected) of the mutation carriers in this study was best categorised by the FDG metabolism in the parietal cortex in 14 of 15 subjects with a sensitivity of 100% and a specificity of 85.7%. As Machado-Joseph disease is a late onset, progressive disorder with preclinical and clinical abnormalities which are age dependent, it is possible that an asymptomatic subject would be normal at some earlier time of life until a certain time point when preclinical changes occur. If enough asymptomatic carriers of younger age were studied, the sensitivity of PET might be considerably lower. The onset of these preclinical changes may be time dependent, occurring in the period just before the onset of symptoms. In other words, the PET changes may inversely correlate with the difference between the age of the asymptomatic subject at the time of PET and the age of onset of symptoms in the affected relative. However, this difference is often difficult to determine. The age at symptom onset may be similar in members of the same generation of the same family. However, there may be considerable variation of ages at onset of symptoms between members of the same family in different generations (anticipation phenomenon), which makes estimation of the difference very unreliable.

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Albrecht von Graefe (1828–70)

Von Graefe, who was of gigantic importance in ophthalmology, died at the early age of 42. By the age of 39 Von Graefe was internationally a unique figure and presided and dominated over the entire 3rd International Congress of Ophthalmology held in Paris in 1867. He read four papers including a classic description of choroid tubercles, but his most notable contribution was his exposition of his “modified linear extraction” as a new technique for the cataract.

His contributions to ophthalmology were multiple. His name is eponymously remembered in the von Graef sign in exophthalmic goitre and the von Graefe extraction knife. Ophthalmology developed through the application of the ophthalmoscope by von Graefe. His clinical contributions included the physiology of the oblique extraocular muscles and the symptoms of ocular paralysis, silver nitrate treatment of conjunctivitis, description of the various types of hemianopia, the efficacy of iridectomy in acute glaucoma, occlusion of the central retinal artery by emboli, the recognition of papilloedema, and the recognition of optic neuritis rather than paralysis of the optic nerve as being a cause of central visual failure.

Von Graefe died from tuberculosis. In 1882 his statue (with the ophthalmoscope) was erected in the garden of Charite Hospital in Berlin, and later relocated to Shuman Avenue.

Along with Donders, von Graefe founded the Archiv für Ophthalmologie, which did so much to raise the status of this specialty. Donders and Arlt became the editors. A stamp was issued by West Germany (West Berlin) in 1978 on the 150th year of the birth of von Graefe. (Stanley Gibbons B553, Scott 9N417).